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HIGHEST RN 929680-66-0 HIGHEST RN 929680-66-0 10 APR 2007 10 APR 2007 STRUCTURE FILE UPDATES: DICTIONARY FILE UPDATES: New CAS Information Use Policies, enter HELP USAGETERMS for details.

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http://www.cas.org/ONLINE/UG/regprops.html

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STRUCTURE UPLOADED . 10

=> d l6 L6 HAS NO ANSWERS L6 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT Structure attributes must be viewed using STN Express query preparation.

=> s 16 SAMPLE SEARCH INITIATED 12:37:39 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLEYED - 384 TO ITEBATE

384 ITERATIONS 100.0% PROCESSED SEARCH TIME: 00.00.01

5 ANSWERS

ONLINE **COMPLETE**
BATCH **COMPLETE** 6505 TO 5 TO PROJECTED ITERATIONS: PROJECTED ANSWERS:

FULL FILE PROJECTIONS:

5 SEA SSS SAM L6

FULL SEARCH INITIATED 12:37:43 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 7528 TO ITERATE => s 16 full

7528 ITERATIONS 100.0% PROCESSED SEARCH TIME: 00.00.01

81 ANSWERS

81 SEA SSS FUL L6 r8

TOTAL SESSION 356.26 SINCE FILE ENTRY 172.10 => file caplus COST IN U.S. DOLLARS

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TOTAL SESSION -0.78 SINCE FILE ENTRY 0.00 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) CA SUBSCRIBER PRICE

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FILE COVERS 1907 - 11 Apr 2007 VOL 146 ISS 16 FILE LAST UPDATED: 10 Apr 2007 (20070410/ED)

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http://www.cas.org/infopolicy.html

=> s 18

23 L8

> s 18 full 23 L8 110 s

=> s 110 py<2003 MISSING OPERATOR L10 PY<2003

The search profile that was entered contains terms or nested terms that are not separated by a logical operator.

=> s 110 and py<2003 22870433 PY<2003 L11 14 L10 AND PY<2003

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Lil ANSWER 1 OF 14 CAPLUS COPYRIGHT 2007 ACS on STN ACESSION NUMBER: 2006:1024194 CAPLUS DOCUMENT NUMBER: 145:397368

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

Preparation of sulfonyl aryl or heteroaryl hydroxamic acid compounds as matrix metalloprotease inhibitors Bedell, Louis J.; Mcdonald, Joseph J.; Barra, Thomas E.; Becker, Daniel P.; Shashidhar, Rao N.; Freskos, John N.; Mischke, Brent V.; Getman, Daniel P.; Decrescenzo, Gary A.; Villamil, Clara I. G. Searle & Co., USA
U.S., 182pp., Cont.-in-part of U.S. Ser. No. 310,813. INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

Patent English 11

DOCUMENT TYPE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. US 2000-569034 US 1999-230209 KIND PATENT NO.

20000511 19990624 <--20010507 <--CN, LLR, PT, US, CH, LK, UG, BZ, GB, KZ, NO, TZ, 71, KR, WO 2001-US14706 WE KE **遇咒强责**克 20061003 20010906 20020430 20011115 20020307 AU, AZ, DK, DM, IN, IS, MD, MG, SI, SK, H SG, B1 B2 B2 A2 A2 C2, ID, IV, SE, SE, EI, ES, YU, YU, US 7115632 US 2001020021 US 6380258 WO 2001085680 WO 2001085680

Erich Leeser

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TD, TG 20010719 19990512 B2 A2 A2 A2 CM, GA, GN, GW, ML, MR, NE, SN, 20030417 US 2001-909227 20040224 US 1999-310813 US 1999-230209 US 1997-35182P WO 1998-US4300 US 2000-569034 US 2000-728408 MARPAT 145:397368 BJ, CF, CG, CI,
US 2003073845 A1
US 6696449 B2
PRIORITY APPLN. INFO.: OTHER SOURCE(S):

ΑB

The title compds. [I; A = O, S, CO2, etc.; R = alkyl, alkoxyalkyl, aryl, etc.; E = CO, SO2, (un)substituted COMH, etc.; Y = H, alkyl, alkoxy, etc.; R5, R6 = H, alkyl, cycloalkyl, etc.; R20 = OR21, NR130R22 etc. (R13 = H, alkyl, benzyl; R21 = alkyl, aryl, arylalkyl; R22 = selectively removable protecting group)! Or pharmaceutically acceptable salts thereof that interallal inhibit matrix metalloprotease activity, are prepared Thus, thoetherification of 4-phenoxybenzenethiol with 2-fluorobenzaldehyde in the presence of R203 in isopropanol under reflux for 20 h gave 2-(4-phenoxyphenylthio)benzaldehyde which was condensed with tetra-Et dimethylaminomethylenediphosphonate in the presence of NaH in THF at room temperature for 4 h gave to 2-[2-(4-phenoxyphenylthio)penyllacetic acid which was condensed with O-tetrahydroxylaminographyllacetic acid which was condensed with O-tetrahydroxylamine using 1-ethyl-2-(3-denoxyphenyllacetic acid which was condensed with O-tetrahydroxylamine using 1-ethyl-2-(3-denoxyphenyllacetic acid which was condensed with Cetrahydroxylaminopropyl) carbodisimide hydrochloride in MeCN followed by treatment with p-toluenesulfonic acid in methanol at room temperature for 2 h

give N-hydroxy-2-[2-(4-phenoxyphenylsulfonyl)phenyl]acetamide (III). III and N-hydroxy-2.3-dimethoxy-6-[[4-[4-(trifluoromethyl)phenoxy]-1-piperidinyl]sulfonyl]benoxide showed IC50 of >10,000 nM against MMP-1, 0.3 and 2.4 nM, resp., against MMP-2, and 2 and 2.7 nM, resp., against MMP-13. Also disclosed is a treatment process that comprises administering a contemplated sulfonyl aromatic or heteroarom. ring hydroxamic acid compound in a matrix metalloprotease (MMP) enzyme-inhibiting effective amount to a host having a condition associated with pathol. MMP activity. RL: PAC (Pharmacological activity): SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES Ľ

to

(preparation of sulfonyl aryl or heteroaryl hydroxamic acid compds. matrix metalloprotease inhibitors)
308385-85-5 CAPLUS 2-[(4-benzoyl-1-piperazinyl)sulfonyl]-N-hydroxy- (9CI)

Z Z

CY, BF,

TR,

BE, SE, AT, PT,

ZW, NL,

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TZ, LU, SZ, IT,

SI, IE,

SB,

RW:

as

308385-86-6 CAPLUS
Benzamide, N-hydroxy-2,3-dimethoxy-6-[[4-[phenylmethyl]-1-piperazinyl]sulfonyl]- (9CI) (CA INDEX NAME) Z Z

RN 308385-87-7 CAPLUS
CN Benzamide, N-hydroxy-2,3-dimethoxy-6-[[4-[[4-([rifluoromethoxy)phenyl]meth
y1]-1-piperazinyl]sulfonyl]- (9CI) (CA INDEX NAME)

THERE ARE 72 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 72 REFERENCE COUNT:

ACCESSION NUMBER:

DOCUMENT NUMBER:

138.304308
TITLE:

PERPARATION of sulfonyl aryl hydroxamates and their use as matrix metalloprotease inhibitors
Barta, Thomas E.; Becker, Daniel P.; Bedell, Louis J.;
Barta, Thomas E.; Becker, Daniel P.; Bedell, Louis J.;
Barta, Thomas E.; Becker, Daniel P.; Bedell, Louis J.;
Barta, Thomas E.; Becker, Daniel P.; Bedell, Louis J.;
Barta, Thomas C.; Jaseph J.; Mischke, Brent V.; Rao, Shashidhar N.; Villamil, Clara I.
PATENT ASSIGNEE(S):
PATENT ASSIGNEE(S):
PATENT ASSIGNEE(S):
Ser. No. 569,034.

Erich Leeser

	DATE	20010719	1000001	9960304		A2,	Ç	CG CT CM	Ì	19990624 <	1110000	20000311	1000	20020719	20020719	CA. CH. CN.	, H	ĽĶ,	ğΊ	TR, TT, TZ,	;	Αζ,	DK, EE, ES, BF, BJ, CF,		20020719	20020719	SE, MC, PT,	SK 20000110	20020719	19970304					20001201	2002071		
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	APPLICATION NO.	22	0001-0001-00		ž Š	UA, US, UZ,		UG, ZW, AT, BE, CH,	(1)	US 1999-230209		US 2000-369034		CA 2002-2453613	WO 2002-US23219	BA. BB. BG. BB. BY.	EC, EE,	KE, KG, KP,	MN, MW, MX,		Z.W.	, T.Z., 2.S.		MR, NE, SN,	AU 2002-32643	EP 2002-76114	GR, IT, LI,		BR 2002-11430	17 2003-313361 115 1997-351829	WO 1998-US4300	US 1999-310813			116 2001-000223			
CODEN: USXXCO Patent English 11	KIND DATE		B2 20040224	٦ د	KP, KR, LC, LK,	SK, SL,	TJ, TM	LS, MW, SD, SZ,	NE, SN, TD,		B2 20020430	B1 20061003				Ę	DE, DK,	IL, IN,	MA, MD,	SD, SE,	Ž,	MM, MZ,	RU, IJ, IM, AI, GR IF, IT, IU.	9N.		20040	DK, ES,	EI,	A 20040/13								MARPAT 138:304308	
UNT:	¥2 i				i i	SG,	Đ.	ž (ğ				•			74	9	HO,	ĽŪ,	80,	us,	¥.	£ 6	ð			E	5		32 TNEO .	:						Σ	
DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. CO	PATENT NO.	200307384	US 6696449		w: AL, AU,		Ϋ́ς:	RW: GH, GM,	GP. GR.	200102002	US 6380258	US /115632	6794511				i 8		LS, LT,		UA,		KG, KZ,		200232643	1406626	R: AT, BE,	IE,		2F ZUUSSUZBSS							OTHER SOURCE(S):	GI

Title compds. I [W = 6-membered heterocycle containing the sulfonyl bonded N; A-R-E-Y = 4-substituent; A = 0, S00-2, etc.; R = alkyl, alkxysalkyl, aryl, heteroraryl, cycloalkyl, etc.; E = absent, bond, C0, S02, etc.; T = absent, H, OH, CN, NO2, alkyl, haloalkyl, aninoalkyl; R5-6 = together with the atoms to which they are bonded, form an aliphatic or aromatic carbocyclic AB ö

heterocyclic ring having 5-7 members) are prepared Over 50 synthetic examples are disclosed. For example, phthalide is reacted with he defibenoxy) benzenethiol (DMF, K2CO3, 100°C, 2 h) and the resulting product converted to the hydroxamic acid (GH2C12, CICOCO1, DMF (cat), TMSOWH2, 0°C, 1.5 h) followed by oxidation (GH2C12, mCPBA, room temperature, 3 h) to II. II has ICS0 = 10 nM for MMP-2, 45 nM for MMP-13 and >10,000 nM for MMP-1. I are inhibitors of MMP and angiogenesis.

308.385-89-58 308385-86-6P 308385-87-7P RE. PAC (Pharmacological activity); SPN (Synthetic preparation); USES (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

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S S

(use of sulfonyl ary) or heteroaryl hydroxamic acids and derivs. as aggrecanase inhibitors)
308382-85-5 CAPLUS
Benzamide, 2-[(4-benzoyl-1-piperazinyl)sulfonyl]-N-hydroxy- (9CI) (CA INDEX NAME)

308385-86-6 CAPLUS Z.

Erich Leeser

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Benzamide, N-hydroxy-2,3-dimethoxy-6-[[4-(phenylmethyl)-1-piperazinyl]sulfonyl}- (9CI) (CA INDEX NAME) Ç

308385-87-7 CAPLUS
Benzamide, N-hydroxy-2,3-dimethoxy-6-[[4-[[4-[trifluoromethoxy)phenyl]meth
yl]-1-piperazinyl]sulfonyl]- (9CI) (CA INDEX NAME) Z Z

COPYRIGHT 2007 ACS on STN ::319307 CAPLUS

L11 ANSWER 3 OF 14 CAPLUS COPYRICH
ACCESSION NUMBER: 1307-19307
DOCUMENT NUMBER: 137:75137
TITLE: Predictions

Professions of Binding of a Diverse Set of Ligands to Gelatinase-A by a Combination of Molecular Dynamics and Continum Solvent Models. Xiaojie College of Chemistry and Molecular Engineering, Peking University, Beijing, 100871, Peop. Rep. China Journal of Physical Chemistry B (2002), 106(21), 5527-5535 CODEN: JPCEFK; ISSN: 1089-5647 AUTHOR(S): CORPORATE SOURCE:

SOURCE:

American Chemical Society Journal DOCUMENT TYPE: LANGUAGE: AB The free en PUBLISHER:

The free energies of binding, Agbind, between a diverse set of eight hydroxamate inhibitors with gelatinase-A (MPM-2) were computed by using the recently developed MVPBSA approach. In this paper, a nonbonded model was used to represent the potentials of the catalytic zinc center. Mole dynamics (MD) simulations were used to generate the thermally averaged ensemble of conformations were used to generate the thermally averaged ensemble of conformations of the ligand-protein complexes. On the basis of the trajectories from MD simulations, the free energies of binding were calculated using mol. mechanics, the continuum solvent model, surface area estimation, and normal-mode anal. The results show that MM/PBSA not only can rank the studied ligands effectively but also can reproduce the exptl. binding free energies successfully. The predicted binding free energies correlate well with the exptl. values (r = 0.84, q = 0.78). As a comparison, the free energies of binding were also computed by using the

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Linear interaction energy approximation (LIE). The overall agreement between the calculated and expell, values for the diverse set of ligands means that the MAMPBA approach is a useful tool for the general evaluation of protein-ligand interactions. The anal. of the sep. energy terms contributing to MAMPBA free energy indicates that the association between hydroxamate and MMP-2 is mainly driven by more favorable van der MMABAS, mainly driven by more favorable van der MMABAS, in mainly driven by more favorable van der

220046-45-7 RL: BSU (Biological study, unclassified); BIOL (Biological study) (linear interaction energy approximation reveals association between

H

hydroxamate

and MMP-2 is promoted by van der Waals/nonpolar interactions in complex than in solution) 220046-45-7 CAPLUS

1-Piperazinebutanamide, α-amino-4-(1,3-benzodioxol-5-ylmethyl)-β-(cyclopentylmethyl)-N-hydroxy-γ-oxo-, (αS,βR)-(9CI) (CA INDEX NAME)

Z Z

Absolute stereochemistry.

THERE ARE 61 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 61 REFERENCE COUNT:

Preparation of hydroxamic acid peptide deformylase inhibitors as antibacterial agents Chong, Lee; Frechette, Roger; Scott, Carole; Tester, Chichard; Smith, Whitney; Chiba, Katsumi; Sakamoto, Masatoshi; Gluchowski, Charles COPYRIGHT 2007 ACS on STN 2002:275960 CAPLUS 136:310184 LIII ANSWER 4 OF 14 CAPLUS ACCESSION NUMBER: 200 DOCUMENT NUMBER: 136 INVENTOR (S):

PATENT ASSIGNEÉ(S):

Questcor Pharmaceuticals, Inc., USA PCT Int. Appl., 171 pp. CODEN: PIXXD2 DOCUMENT TYPE:

Patent English FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

20010924 <--CA, CH, DATE , ZE BR, BY, WO 2001-US29926 APPLICATION NO. BG, BA, BB, 20020411 AT, AU, AZ, DATE KIND 22 AE, AG, AL, WO 2002028829 WO 2002028829 PATENT NO.

Erich Leeser

я, Рг, Уб, 8,8,8,8 ЯК, Ч**Я**, 8 8 K GD, NZ, TZ, 3 E 8 KZ, TT, GE, 73, X3, ZW, ES, 2000-234967P 2001-761850 2001-US29926 KP, 유. 유. KG, MW, 7Z, DE, BJ, SE, KE, SZ, CY, BF, AU US WO SI, TR, MARPAT 136:310184 MZ, SD, AT, BE, PT, SE, SN, TD, 20020415 DM, IS, MG, SI, S, E, E, SD, ZA, IS, TJ, MC, AS, KE BE AU 2002030385 PRIORITY APPLN. INFO. GW. 5 £ £ 5 2002030385 OTHER SOURCE(S): GI RW:

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Hydroxamic acid derivs. of peptides and peptidomimetics of formulas I, II, and III (wherein Z = NHOH or OR3: Ra = alkyl or a biocleavable moiety; X = and III (wherein Z = NHOH or OR3: Ra = alkyl or a biocleavable moiety; X = (CO or \$02; Y = (un)substituted heteroalkyl or heterocycly; RS3 = (un)substituted (cyclolaikyl, aryl, beterocycly, or heteroalkyl; RZR3 = 4-7 membered (un)substituted heterocycle; RZR4 = ring formed through a CH2CH2 linkage; or R2 = Me; or R3 = Hor (un)substituted (hetero)alkyl, aryl, or heterocyclyl; R5 and R6 = independently H, NOZ, NH2, NHCOH, NHCOCH3, NHSOCH3, or (un)substituted GRNH-(hetero)alkyl or CHZNH-heterocyclyl; one of R7 or R8 = CHR10COMHOH; one of R7 or R8 = (un)substituted (hetero)alkyl, alkyl) heterocyclyl, or alkylaryl; R9 and R10 = independently H or (un)substituted (hetero)alkyl, (alkyl)heterocyclyl, or alkylaryl; R9 and R10 = alkylaryl) were prepared as peptide deformylase (Fe-PB) inhibitors for alkylaryl) were prepared as peptide deformylase (Fe-PB) inhibitors for treating various bacterial infections. For example, 3-pyrrolidinal massing treating various processes and the second secon reading various bacterial infections. For example, 3-pyrrolidinol was added to tert-Bu (R)-(2-pentyl) succionate mono(R-Hydroxysuccinninde) estet to give the amide (68). Treatment with 20% TFA/DCM, followed by MeOH, benzene, and TMSN2 in hexanes, to afford the Me ester (90%). The pyrrolidinol was coupled with 4-methoxyphenylisocyamate and the ester converted to the hydroxamic acid (IV) using NH2OH+HCl. The latter inhibited E. coli Fe-PDF with IC50 of 9 nM and showed selectivity for Fe-PDF vs. thermolysin with a selectivity index of 30,000. Thus, I, II and II are useful as antibiotics against a broad range of infectious Æ

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES 409129-95-9P 409129-96-0P Uses) H

(peptide deformylase inhibitor; preparation of hydroxamic acid derivs. of peptides and peptidomimetics as peptide deformylase inhibitors for treatment of infectious diseases) 409129-95-9 CAPLUS

Z Z

1-Piperazinebutanamide, 4-benzoyl-N-hydroxy-γ-oxo-β-pentyl-, (βR) - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

409129-96-0 CAPLUS 1-Piperazinebutanamide, N-hydroxy-r-oxo-B-pentyl-4-(1-pyrrolidinylcarbonyl)-, (BR)- (9CI) (CA INDEX NAME) £ 5

Absolute stereochemistry.

Hou, Ting-Jun; Zhang, Wel; Xu, Xiao-Jie College of Chemistry and Molecular Engineering, Peking University, Beljing, 100871, Peop. Rep. China Huaxue Xuebao (2002), 60(2), 221-227 CODEN: HHHPA4; ISSN: 0567-7351 137:5788 Binding free energy calculations for MMP2-hydroxamate LII ANSWER 5 OF 14 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:161702 CAPLUS DOCUMENT NUMBER: 137:5788 Binding free energy calculation Kexue Chubanshe complexes AUTHOR(S): CORPORATE SOURCE: PUBLISHER: DOCUMENT TYPE: LANGUAGE: AB The absolu SOURCE:

The absolute binding affinities of hydroxamate inhibitors with MMP-2 were evaluated by mol. dynamics (MD) simulations with a linear response evaluated by mol. dynamics (MD) simulations with a linear response approach. During MD simulations, a nonbonded model for the catalytic Zn center was used to represent the interactions between Zn center and enzyme/inhibitor. The trajectories from MD simulation show that using the nonbonded model the catalytic Zn ion adopts five coordination number, but the coordination form exists large difference with that of the initial model. After fittings, the models with one parameter, two parameters and three parameter model with a constant term bears the best predicting ability. The best model with a constant term bears the best predicting ability. The best model yields an average error of 2.38 kJ/mol for the eight producting affinities of hydroxamates. H

RL: BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study) (binding free energy calcns. for MMP2-hydroxamate complexes) 220046-45-7 Aprus 220046-45-7

220046-45-7 Z

Erich Leeser

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1-Piperazinebutanamide, α-amino-4-(1,3-benzodioxol-5-ylmethyl)-β-(cyclopentylmethyl)-N-hydroxy-γ-oxo-, (αS,βR)-(9CI) (CA INDEX NAME) Z

Absolute stereochemistry.

Beckel, Louis J.; Mconald, Joseph; Barta, Thomas E.; Becker, Daniel P.; Rao, Shashidhar N.; Freskos, John N.; Mischke, Brent V.; Getman, Daniel P.; Decrescenzo, Gary A.; Villamil, Clara I. Pharmacia Corporation, USA PCT Int. Appl., 374 pp. Preparation of sulfonyl aryl or heteroaryl hydroxamic acid compounds as inhibitors of matrix WO 2001-US14706 APPLICATION NO. L11 ANSWER 6 OF 14 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2001:833270 CAPLUS DOCUMENT NUMBER: 135:371526 TITLE: Preparation of wellows metalloproteinase 20011115 Patent English 11 KIND LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT ASSIGNEE(S): WO 2001085680 WO 2001085680 PATENT NO. DOCUMENT TYPE: INVENTOR(S): SOURCE:

20010507 <--A 20000511 B2 19990512 A2 19990624 CH, CY, TR, BF, 20000511 SE, 4,2,5,4 4,7,5,4 PT, TD, BZ, GB, KZ, TZ, MZ, TT, ZW, NL, SN, US 2000-569034 US 2000-569034 US 1999-310813 US 1999-230209 RP, TR, MC, M. KG, 12, 10, MR, S2, IT, ML, BA, JP, JP, KK, SE, MARPAT 135:371526 MZ, SD, GB, GR, GA, GN, 20061003 AZ, DM, IS, MG, SK, SI, YE, A2 A3 AM, CZ, IIV, IV, SE, ZA, ILS, CI, CI, US 7115632 PRIORITY APPLN. INFO.: AE, AG, GM, HR, LS, LT, RO, RU, UZ, VN, GH, GM, BJ, CF, OTHER SOURCE(S): GI RW:

AB Title compds. I [W = 5-, 6-membered aromatic or heteroarom. ring; R1 = a substituent containing a 5- or 6-membered cyclohydrocarbyl, heterocyclo, arylor betacrary radical that is bonded directly to the depicted SO2-group or heteroaryl radical that is bonded directly to the depicted SO2-group said R1 with certain steric requirements; R5-6 = H, alkyl, cycloalkyl, acylalkyl, halo, nitro, hydroxy, cyano, alkoxy, haloalkyl, haloalkyloxy, hydroxylkyl, etc. or R5-6 fogether with the atoms to which they are bonded form a further aliphatic or aromatic carbocyclic or neterocyclic ring having 5-to 7-members; R20 = 0R21, where R21 = H, alkyl, aryl, arylalkyl, NR130R22, where R22 = a selectively removable protecting group and R13 = H, alkyl, benzyl group, etc.] were prepared Over 50 synthetic examples were disclosed. For example, phthalide was reacted with 4-tomeonethiol (DMF, R2CO3, 100°C, 2 h) and the resulting product converted to the hydroxanic acid (CH2C12, CLCOCC1, DMF (cat), TMSONH2, 0°C, 1.5 h) followed by oxidation (CH2C12, CLCOCC1, DMF (cat), TMSONH2, 0°C, 1.5 h) followed by oxidation (CH2C12, CLCOCC1, almost and 213 and 210,000 and for MMP-1. I are inhibitors of MMP and angiogenesis.

IT 30835-85-5p, 2-[(4-Benzoyl-1-piperazinyl)sulfonyl)-N-hydroxyle-[(4-hydroxyl-2,3-dimethoxy-6-[(4-hydroxyl-2,3-dimethoxyl-6-[(4-hydroxyl-2,3-dimethoxy æ

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Blological study); PREP (Preparation); USES (Uses) (drug; preparation of sulfonly aryl or heteroaryl hydroxamic acid compds. as inhibitors of matrix metalloproteinase) (trifluoromethoxy)phenyl]methyl]-1-piperazinyl]sulfonyl]benzamide hydrochloride II

<u>g</u> 2-[(4-benzoyl-1-piperazinyl)sulfonyl]-N-hydroxy- (9CI) 308385-85-5 Benzamide, 2 INDEX NAME ₹ 3

Erich Leeser

50613257

373367-17-0 CAPLUS
Benzamide, N-hydroxy-2,3-dimethoxy-6-[[4-(phenylmethyl)-1piperazinyl]sulfonyl]-, monohydrochloride (9CI) (CA INDEX NAME) Z Z

● HC1

Benzamide, N-hydroxy-2,3-dimethoxy-6-[[4-[[4-(trifluoromethoxy)phenyl]methyl]-1-piperazinyl]sulfonyl]-, monohydrochloride (9CI) (CA INDEX NAME) 373367-18-1 CAPLUS Z Z

L11 ANSWER 7 OF 14 CAPLUS COPYRIGHT 2007 ACS on STN ACESSION NUMBER: 2001:472692 CAPLUS DOCUMENT NUMBER: 135:61355 ritle:

Preparation of α-arylethylpiperazine derivatives as neurokinin antagonists Stiernet, Francoise; Genicot, Christophe; Lassoie, Marte-agnes; Moureau, Florence; Ryckmans, Thomas;

INVENTOR (S):

Taverne, Thierry; Henichart, Jean-pierre; Neuwels, Michel; Goldstein, Solo Ucb, S.A., Belg. PCT Int. Appl., 115 pp. CODEN: PIXXD2 Patent English DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COI PATENT INFORMATION: PATENT ASSIGNEE(S): SOURCE:

COUNT:

KIND ----Al AM, A' DE, D ES, PE, 도, B. A. T A1 B2 SI, SI, W: AE, AG, AL, A CR, CU, CZ, D LU, LV, MA, M SU, SE, SG, S YU, ZA, ZW, S RW: GH, GM, KE, L DE, DK, ES, F BJ, CF, CG, C д**,** Ë, JP 2003518108 US 2003220323 US 6916797 PRIORITY APPLN. INFO.: R: AT, BE, IE, SI, R: AT, BE, IE, SI, WO 2001046167 OTHER SOURCE(S): GI EP 1242399 PATENT NO.

MARPAT 135:61355

The title compds. [I: Z = O, S: nl = 1-2; R2 = H, Me; W = cyclohexyl substituted by a COSH, 2-phenylacetic acid, or alkyl 2-phenylacetate, etc.; Axl = (un)substituted Ph, aryl, heteroaryl, etc.; Axl = (un)substituted Ph, etc.] and their salts, useful as neurokinin receptor antagonists (NKlantagonists), were prepared Thus, hydrolysis of the corresponding Et ester afforded [[2 = O, R2 = H, nl = 1; W = (CH2) 4CO2H, Axl = Ph, Rx2 = 3,5-(F30) 2G6H3) which showed piC50 of 7.5 against binding to NKl receptors. The compds. I are useful for the prevention and/or treatment of a condition associated with pathol. levels of substance P. Ris. BAC (Blological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); ΑB H

Erich Leeser

BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of $\alpha\text{-arylethylpiperazine derivs.}$ as neurokinin

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antagonists)
36416-43-1 CAPLUS
1-Piperazinehexanamide, 4-[2-[[3,5-bis(trifluoromethyl)phenyl]methoxy]-1-phenylethyl]-N-hydroxy- (9CI) (CA INDEX NAME) S S

HO-NH-C- (CH2) 5

20001214 <--

2000-EP12667

20010628

BB, ES,

APPLICATION NO.

346416-44-2 CAPLUS 1-Piperazinehexanamide, 4-[2-[[3,5-bis(trifluoromethyl]phenyl]methoxy]-1-phenylethyl]-N-hydroxy-, (22)-2-butenedioate (1:2) (salt) (9CI) (CA INDEX NAME) S S

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19991220 <--

NL, SE, MC, PT,

BE, CH, CY, SE, TR, BF, TG

AT, PT, TD,

SZ, TZ, UG, ZW, A, IT, LU, MC, NL, P, ML, MR, NE, SN, T EP 1999-125359

LR, LS, PT, RO, US, UZ,

BY, IC, IC, UA, UA, SW,

MZ, TT, TZ, LU, MR, MX, MD, SZ, SZ, III, ML, ML,

SE, SE, GW,

BZ, GE, LK, PL,

20001214 <--NL, SE, MC, PT,

20001214 20020830

3, GR, IT, LI, LU, N , AL, TR JP 2001-547078 US 2002-168331

GB,

ES, FR, RO, MK,

DK,

EP 2000-989974

GB,

A 19991220 W 20001214

EP 1999-125359 WO 2000-EP12667

346416-43-1 C27 H33 F6 N3 O3 CRN

HO-NH-C- (CH2) 5

7 S CRN 110-16-7 CMF C4 H4 O4

Double bond geometry as shown.

CO2H

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT REFERENCE COUNT:

L11 ANSWER 8 OF 14 CACCESSION NUMBER:
DOCUMENT NUMBER:
TITLE:

CAPLUS COPYRIGHT 2007 ACS on STN 2001:390470 CAPLUS 135:10410 SAFIN SAFI

with a Linear Interaction Energy Approach Hou, T. J., Zhang, W., Xu, X. J. College, C. Chemistry and Molecular Engineering, Peking University, Beijing, 100871, Peop. Rep. China Journal of Physical Chemistry B (2001), 105(22), 5304-5315 CODEN: 1089-5647 AUTHOR(S): CORPORATE SOURCE:

SOURCE:

American Chemical Society Journal

English

PUBLISHER: DOCUMENT TYPE: LANGUAGE: AB The bindin

Ab The binding of a series of hydroxamate inhibitors with gelatinase-A is examined to evaluate the viability of calculating free energies of binding, AGD, utilizing mol. dynamics (MD) simulations with a linear interaction energy approach. In our simulations, a bonded model was used to represent the potentials of the catalytic zinc center. The electrostatic distribution of this model was derived using a two-stage electrostatic distribution of this model was derived using a two-stage electrostatic potential fitting calcus. The resulting bonded model was the nergies of binding for the 15 hydroxamate inhibitors. In the correlation process, several linear models consisted of different energy components mergies of binding for the 15 hydroxamate inhibitors. In the correlation process, several linear models consisted of different energy components were tested. We found that besides the usually used Coulombic and var der wasts energy terms, the introduction of a constant term could significantly improve the correlation. The best model yields an average error of 0.6 kcal/mol. The predictive ability of the best model was revealed by the high value of q2 (0.854) from the leave-one-out cross-validation. To this series of inhibitors, and also provided insights into the interactions occurring in the active site and the origins of variations in AGD. The PI' groups of inhibitors and energies with the ninearcing studied inhibitors and energies and several model was revealed by the high simulations predicted the binding mode of the gelatinase-A with the scrive site and the origins of variations in AGD. The PI' groups of inhibitors make extensive van der Waals and Mydrophobic contacts with the nonpolar side chains of four residues in AGD. The PI' groups of inhibitors make extensive van der Waals and Mydrophobic contacts with the nonpolar side chains between hydroxamates and elatinase-A are very important to stabilize the inhibitors in the active site the hydrogen bonds between the P3' group and gelatinase-A can produce more favora

unclassified); PRP RI: BPR (Biological process); BSU (Biological study, unclassified); R (Properties); BIOL (Biological study); BROC (Process) (binding affinities of a series of selective inhibitors of gelatinase-A using mol. dynamics with a linear interaction energy

H

approach) 220046-45-7 Z Z

1-Piperazinebutanamide, α-amino-4-(1,3-benzodioxol-5-ylmethyl)-β-(gyclopentylmethyl)-N-hydroxy-γ-οxo-, (αS,βR)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

Erich Leeser

50613257

THERE ARE 52 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 25 REFERENCE COUNT:

Pharmacia, Department of Medicinal Chemistry, Skokie, 11., 6007, USA Bioorganic & Medicinal Chemistry Letters (2000), 10(04), 2815-2817 CODEN: BMCLES; ISSN: 0960-894X Elsevier Science Ltd. Barta, T. E., Becker, D. P.; Bedell, L. J.; De Crescenzo, G. A.; McDonald, J. J.; Munie, G. E.; Rao, S.; Shieh, H.-S.; Stegeman, R.; Stevens, A. M.; Villamil, C. I. Synthesis and activity of selective MMP inhibitors US COPYRIGHT 2007 ACS on STN 2000:853658 CAPLUS with an aryl backbone CASREACT 134:222499 134:222499 Journal CAPLUS A series of novel, L11 ANSWER 9 OF 14 ACCESSION NUMBER: DOCUMENT NUMBER: DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): CORPORATE SOURCE: AUTHOR(S): PUBLISHER: SOURCE:

N-hydroxy-2-[[4-(phenylmethyl)-1-piperidinyl]sulfonyl]benzamide,
2-fluoro-N-hydroxy-6-[[4-(4-(trifluoromethyl)phenoxy]-1piperidinyl]sulfonyl]benzamide, and derivs. or homologs thereof. The
crystal and mol. structure of 2-fluoro-N-hydroxy-6-[[4-(4-(4-(trifluoromethyl)phenoxy]-1-piperidinyl]sulfonyl]benzamide compound with MMP-1 sparing arylhydroxamate sulfonamides with IP-2 and MMP-13 is described. Example compds. tl tested were N-hydroxy-2-[[(phenylmethyl)amino]sulfonyl]benzamide, N-hydroxy-2-[[(4-methoxyphenyl)methylamino]sulfonyl]benzamide activity against MMP-2

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study) ((annocultonyl)-N-hydroxybenzamide derivs. and their activity as gelatinase (MMP-2) and collagenase (MMP-13) inhibitors) MMP-8 were reported. 308385-85-5 H

S 23

Į 2-[(4-benzoyl-1-piperazinyl)sulfonyl]-N-hydroxy- (9CI) INDEX NAME

THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT . REFERENCE COUNT:

Preparation of hydroxamic acid derivatives as matrix metalloprotease inhibitors
Bedall, Louis J.; McDonald, Joseph J.; Barta, Thomas E.; Becker, Daniel P.; Rao, Shashidhar N.; Freskos, John N.; Mischke, Brent V.; Getman, Daniel P.; G.D. Searle and Co., USA
PCT Int. Appl., 380 pp.
PCT Int. Appl., 380 pp.
Patent
English CAPLUS COPYRIGHT 2007 ACS on STN 2000:824218 CAPLUS LII ANSWER 10 OF 14 ACCESSION NUMBER: DOCUMENT NUMBER: PATENT, ASSIGNEE(S): SOURCE: INVENTOR (S):

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: DOCUMENT TYPE:

CH, CN, CR, GM, HN, HU, LS, LT, LU, SD, SE, WN, YU, ZA, ZW CH, CY, DE, BF, BJ, CF, 20000512 <--20000512 <--SE, MC, PT, 20000512 <--20000512 <--20000512 20000512 20011031 19990512 20000512 20000512 **43** Ĭ, MR, NE, SN, TD, TG 3 CA 2000-2373500 6 EP 2000-931910 7, GB, GR, IT, LI, LU, N BR 2000-11291 JP 2000-618236 NZ 2000-515197 AU 2000-49718 ZA 2001-9007 US 1999-310813 WO 2000-US6713 APPLICATION NO. õ W. SD, WE, SD, AT, KIND St., G., T.S., 5, B A B ¥, ₩ A1 A F 4 A Ξť, M. AE, AE, AL,

W. AE, AE, AL,

CO, C2, DE,

ID, IN, MA, MD,

SG, SI, SK,

RW: GH, GK, ES, FI,

CA 2373500

EP 117773

R. TAT,

BR 2000011291

JP 2002544257

NZ 515197

AU 781399

ZA 201009007

PRIORITY APPLN. INFO.: WO 2000069819 PATENT NO.

Erich Leeser

MARPAT 134:4752

OTHER SOURCE(S): GI

50613257

11

membered cyclohydrocarbyl, heterocyclo, aryl, heteroaryl; RS. R6 independently = hydrido, alkyl, cycloalkyl, acylalkyl, halo, nitro, alkoxyl, acylalkyl, bydroxyalkyl, etc. R20 = alkoxyl, cyano alkoxyl, haloalkyloxy, hydroxyalkyl, etc. R20 = acyptable salts with inter alia inhibite matrix metalloprotease activity are disclosed and a treatment that comprises administering a contemplated sulfonyl aromatic or heteroarom. Hydroxamic acid in an MMP enzyme-inhibiting effective amount to a host having a condition associated with pathol. matrix metalloprotease activity are claimed. Thus, the title compound II was prepared and MMP-2, MMP-3, MMP-13, and MT-MMP inhibition activities Title compds. [1; W = 5, 6 membered aromatic, heteroarom. ring; R = 5, 6 membered cyclohydrocarbyl, heterocyclo, aryl, heteroaryl; R5. R6 ЯВ

were assayed.
308385-85-85-86-6P 308385-87-7P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of hydroxamic acid derivs. as matrix metalloprotease Ħ

308385-85-5 CAPLUS Benzamide, 2-[(4-benzoyl-1-piperazinyl)sulfonyl]-N-hydroxy- (9CI) inhibitors) S S

INDEX NAME)

õ

308385-86-6 CAPLUS
Benzamide, N-hydroxy-2,3-dimethoxy-6-[[4-(phenylmethyl)-1-piperazinyl]sulfonyl]- (9CI) (CA INDEX NAME) C. R.

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Benzamide, N-hydroxy-2,3-dimethoxy-6-[[4-[[4-[trifluoromethoxy)phenyl]methy]]-1-piperazinyl]sulfonyl]- (9Cl) (CA INDEX NAME) 308385-87-7 CAPLUS Z Z

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT REFERENCE COUNT:

Preparation of amino acid sulfonamide hydroxamates Inhibitors of procollagen C-proteinase.
Billedeau, Roland Joseph; Broka, Chris Allen;
Campbell, Jeffrey Allen; Chen, Jian Jeffrey;
Dankwardt, Sharon Marie; Delaet, Nancy; Robinson,
Leslie Ann; Walker, Keith Adrian Murray
F. Hoffmann-La Roche A.-G., Switz.
CODEN: PIXXD2 CAPLUS COPYRIGHT 2007 ACS on STN 2000:441768 CAPLUS 133:74324 Patent English LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: LII ANSWER 11 OF 14
ACCESSION NUMBER:
DOCUMENT NUMBER:
TITLE: PATENT ASSIGNEE (S): DOCUMENT TYPE: INVENTOR (S):

19991214 <--19991214 <--CN, CU, CZ, IL, IN, IS, MA, MD, MG, SI, SK, SL, CH, CY, DE, BF, BJ, CF, SE, TZ, UG, ZW, AT, II, LU, MC, NL, PT, 1, NE, SN, TD, TG CA 1999-2355902 BR 1999-16504 SE, E, E, APPLICATION NO. BG, GH, YU, YU, TZ, LU, NE, AU, FI, A A PIS, TA, GER, ER, 48844889 WO 2000037436 M: AE, I DE, I DE, I TJ, RW: GH, CA 2355902 BR 9916504 PATENT NO.

Erich Leeser

19991214 19991222 <---20010619 <---20010620 20010621 <---20010620 P 19981222 P 19990803 P 19991108 W 19991214 A3 19991222 GR, IT, LI, LU, NL, SE, MC, PT, 20021009 US 1998-113311P US 1999-147053P US 1999-164138P WO 1999-EP9920 US 1999-469660 2001-CN859 2001-3100 US 2002-267727 ZA IN NO US 20040630 ES, FR, GB, C 20031023 20050118 20031120 20040907 H, A1 DE, LV, 172, A2 A2 A1 A1 A1 A1 A1 A1 B2 B1 B2 B1 Ħ, PRIORITY APPLN. INFO.: R: AT, BE, IE, SI, TR 200101868 HU 200104658 JP 200253322 AU 769319 NZ 512292 AU 20271 RU 23271 US 6492394 HR 2001000443 ZA 2001005014 IN 2001CN00859 NO 2001093100 US 6844366 US 2003216405 US 6787559 US 5787559 EP 1149072 EP 1149072

HOHNCOCIRINRSOZARZ [R1 = altyl, haloalkyl, heteroalkyl, cycloalkyl, aryl, aralkyl, aralkenyl, heteroaryl, heteroaralkyl, aminl, aryl, aralkyl, etc.; R = CHRZARL, CHRZELICHARI, RAZ = specified (substituted) Bh, naphthyl; RZ = H, alkyl; with provisos), were prepared Thus, N-hydroxy-2(R1-[(3,4-methylenedioxybenzyl) (4-methoxy-2,3,6-trimethylbenzenesulfonyl) amino]-3-methylbutyzanide was prepared by solution phase synthesis from BOC-D-Val-OH. Title compds. inhibited procollagen C-proteinase with 1550 0.01-2 \muM. 279255-56-0P 279255-58-2P RL: BAC (Biological activity or effector, except adverse); BSU (Biological study), unclassified); SPN (Synthetic preparation); TMU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (USES) MARPAT 133:74324 OTHER SOURCE(S):
AB HOHNCOCHRINRSOZAr2

H

1-Piperazinepentanamide, α -[(1,3-benzodioxol-5-ylmethyl)[(4-methoxyphenyl)sulfonyl]amino]-4-benzoyl-N-hydroxy- δ -oxo-, (αR) - (9CI) (CA INDEX NAME) procollagen C-proteinase) 279255-56-0 CAPLUS S S

Absolute stereochemistry

279255-58-2 CAPLUS
1-Piperazinepentanamide, α-[(1,3-benzodioxol-5-ylmethyl)[(4-methoxyphenyl)sulfonyllaminol-4-(2-furanylcarbonyl)-N-hydroxy- δ-oxo-(αR)- (9CI) (CA INDEX NAME) Z Z

THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 11 REFERENCE COUNT:

Preparation of arylpiperazines as metalloproteinase inhibiting agents (MMP)
Barlant Bernard Christophe; Newcombe, Nicholas John;
Tucker, Howard; Waterson, David
Zeneca Limited, UK; Zeneca-Pharma Sa
PCT Int. Appl., 82 pp. CAPLUS COPYRIGHT 2007 ACS on STN 2000:161258 CAPLUS L11 ANSWER 12 OF 14 C ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: PATENT ASSIGNEE (S): INVENTOR(S): SOURCE:

English DOCUMENT TYPE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

19990825 <---19990825 <--19990825 <--CN, CR, CU, HU, ID, IL, LU, LV, MD, SG, SI, SK, CY, DE, DK, BJ, CF, CG, SE, MC, PT, GB, GR, IT, LI, LU, NL, CH, HR, LT, SE, CH, BY, CA, GH, GM, LR, LS, RU, SD, YU, ZA, AT, BE, TG, SE, APPLICATION NO 20010522 20010627 20060517 DK, ES, FR, C FI, RO, CY AU, KG, AI DM, KE, KE, KE, KE, GN, GS, AI AI AI AI AI AI ******* н, Н, SI, WO 2000012478 44.8.4.5.9.F.9. CZ, IN, SI, SI, CI, CA 2339761 AU 9955247 AU 764367 BR 9913255 EP 1109787 PATENT NO. R.

Erich Leeser

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TR 200100605	T2	20010821	TR	2001-200100605		19990825	;
HU 200103344	A2	20020228	H	2001-3344		19990825	;
EE 200100106	ø	20020617	띮	2001-106		19990825	;
JP 2002523493	F	20020730	Ъ	2000-567511		19990825	;
NZ 509730	æ	20030530	NZ	1999-509730		19990825	
. RU 2220967	C5	20040110	æ	2001-108591		19990825	
NZ 524921	æ	20041029	NZ	1999-524921		19990825	
	Ľ	20060615	. AT	1999-941751		19990825	
	۴	20060929	占	1999-941751		19990825	
	13	20061201	S	1999-941751		19990825	
TW 240722	В	20051001	ž	1999-88114833		19990830	
ZA 2001001231	A	20020513	ZA	2001-1231		20010213	;
US 6734184	B1	20040511	S	2001-763709		20010226	
	ď	20010425	8	2001-1023		20010228	ļ
NO 321478	B1	20060515					
	Æ	20011231	BG	2001-105369		20010322	ţ
HK 1036060	A1	20061027	ΉK	2001-106732		20010924	
AU 2003262101	A1	20031218	AU	2003-262101		20031112	
US 2004171641	Al	20040902	Sn	2004-78775		20040226	
PRIORITY APPLN. INFO.:			ם	1998-402144	ø	19980831	
			ΕP	1999-401351	ď	19990604	
			ş	1999-GB2801	3	19990825	
			ns	2001-763709	A1	20010226	
OTHER SOURCE(S): GI	MARPAT	MARPAT 132:207849					

H, halo, NOZ. etc.; n = 1-3; P = (CH2)n (wherein n = 0-2), alkene, alkyne, etc.; A = (un) substituted 5-7 membered alphatic ring; XI, XZ = N, C, where a ring substituent on ring A is a oxo group that is preferably adjacent a ring N atom; Y = SO2, CO; Z = CONHOH, Y = CO and Q = CR67, CR6R7CH2, NR6, NR6ARZ (wherein R6 = H, alkyl, aralkyl, etc.; R7 = H, alkyl, R f Cogether with R6 forms a carbocyclic or heterocyclic spiro 5-7 membered ring, the latter containing at least one heteroatom selected from N, O, S); Z = CONHOH, Y = SO2 and Q = CR6R7, CR6RCH2, Z = N(OH)GHO, and Q = CR6R6, CR6CH2, NR6GH2; R1 = H, alkyl, cycloalkyl, etc.; R2 = H, alkyl, aryl, etc.], useful as metalloproteinase inhibitors (no data), especially as inhibitors of The title compds. [I, B = monocyclic or bicyclic alkyl, aryl, etc.; R3 = AB

MMP 13, in treating arthritis and atherosclerosis, were prepared E.g., a multi-step synthesis of the title piperazine II was given. Compds. I are effective at 0.5-30 mg/kg/day. H

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of arylpiperazines as metalloproteinase inhibiting agents

260438-45-7 CAPLUS Propananide, N-hydroxy-3-[[4-(phenylmethyl)-1-piperazinyl]sulfonyl]-INDEX NAME)

Z 2

HO-NH-G-CH2-CH2-

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

REFERENCE COUNT:

Preparation of succinyl piperidinamides, morpholinamides, piperazinamides, and analogs as matrix metalloproteinase inhibitors Alpegiani, Marsolino, Pierluigi; Abrate, Francesca; Perrone, Ettore; Corigli, Riccardo; Jabes, Pharmacia & Upjohn S.P.A., Italy PCT Int. Appl., 81 pp.. CODEN: PIXXD2 COPYRIGHT 2007 ACS on STN 64787 CAPLUS 1999:64787 130:139360 English 1 Daniela Patent CAPLUS LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: L11 ANSWER 13 OF 14 OF ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: INVENTOR(S):

19980707 <--19980707 <--ID, IL, JP, KR, MX, NO, NZ, PL, RO, MD, RU, TJ, TM FI, FR, GB, GR, IE, IT, LU, MC, NL, 19970710 19971118 19980707 19980707 19980707 19990310 **443** CA 1998-2265671 AU 1998-88583 EP 1998-940170 JP 1999-508146 US 1999-147798 GB 1997-14548 GB 1997-24395 WO 1998-EP4220 APPLICATION NO. WO 1998-EP4220 CN, CZ, HU, BY, KG, KZ, I DE, DK, ES, 19990208 IT, SE 20010116 20021119 19990121 19990123 KIND A1 A1 A1 B1 B1 G.₹.9 Ë R: DE, ES, F]
JP 2001500533
US 6482827
PRIORITY APPLN. INFO.: AU, BE, C W: AL, I UA, I RW: AT, I CA 2265671 AU 9888583 EP 925289 PATENT NO. WO 9902510

Erich Leeser

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MARPAT 130:139360 OTHER SOURCE(S): GI

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Title compds. I [W = CONHOH or COOH; R1 and R2 = H or an organic residue; R3 = organic group; Q = secondary or tertiary acyclic or cyclic anido group] and their pharmaceutically acceptable salts, solvates, and hydrates are disclosed as inhibitors of matrix metalloproteinases (MMPS), and of the release of tumor necrosis factor-alpha (TNF) from cells. The compds. are therefore useful in the prevention, control and treatment of diseases in which MMPs or TNF are involved, especially tunoral and inflammatory diseases. Processes for their preparation, and pharmaceutical compns. containing them as also described. For instance, the intermediate (45)-(benzyloxycarbonyl)-1-(tert-butoxycarbonyl)-3(R)-isobutylazetidin-2-one (II; preparation given) was subjected to a sequence of ring opening/amidation with piperidine followed by Mydrogenolytic deprotection of the benzyl ester, anidation with PhCHZONHZ HCJ, another hydrogenolysis of the benzyl ester, and acidic deprotection of the BOC-amino group, to give title compound III. The latter compound showed superior aqueous solubility (> 95 mg/mL at 25°), and had Ki values as follows: MMP-1 0.088, MMP-2 0.29, and MMP-3 2.5, all in µM. ЯВ Ħ

RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); PROC (Process); RACT (Reactant or reagent); USES (Uses)

(target compound; preparation of succinyl piperidinamides, morpholinamides, and piperazinamides as matrix metalloproteinase inhibitors) 220046-45-7 CAPLUS 1-Piperazinebutanamide, α-amino-4-(1,3-benzodioxol-5-ylmethyl)-β-(cyclopentylmethyl)-N-hydroxy-γ-oxo-, (αS,βR)-

Absolute stereochemistry

3-(cyclopentylmethyl)-| (9CI) (CA INDEX NAME)

Z Z

H

220046-44-6P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (target compound; preparation of succinyl piperidinamides, morpholinamides, and piperazinamides as matrix metalloproteinase inhibitors)

S S

220046-44-6 CAPLUS carbantc acid, [15,2R)-3-[4-(1,3-benzodioxol-5-ylmethyl]-1-piperazinyl]-2-(cyclopentylmethyl)-1-[(hydroxyamino)carbonyl]-3-oxopropyl]-, 1,1-dimethylethyl ester (9C1) (CA INDEX NAME)

Absolute stereochemistry.

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Z Z

1-Piperazinebutanamide, 4-(1,3-benzodioxol-5-ylmethyl)- β -(cyclopentylmethyl)- α -(dimethylamino)-N-hydroxy- γ -oxo-, (α S, β R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry

Erich Leeser

50613257

220046-57-1 CAPLUS
1-Piperazinebuteanande, 4-(1,3-benzodioxol-5-ylmethyl)-β(cyclopentylmethyl)-N-hydroxy-α-[[(4-methoxyphenyl)sulfonyl]amino]γ-οxo-, (αS,βR)- (9CI) (CA INDEX NAME) S S

Absolute stereochemistry.

CAPLUS 220046-70-8

1-fiperazinebutanamide, α-amino-4-(1,3-benzodioxol-5-ylmethyl)-N-hydroxy-γ-oxo-β-(3-phenylpropyl)-, (ας,βR)-, mono(trifluoroacetate) (salt) (9CI) (CA INDEX NAME) S 2

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CRN 220046-69-5 CMF C25 H32 N4 O5

Absolute stereochemistry,

7 ð CRN 76-05-1 CMF C2 H F3 O2

F- C- CO2H

Z Z

220046-82-2 CAPLUS
1-Pipcerazinebutanamide, α-amino-4-(1,3-benzodioxol-5-ylmethyl)-Picyclopentylmethyl-N-hydroxy-γ-οxo-, (ας,βR)-, bis(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

. G

CRN 220046-45-7 CMF C22 H32 N4 O5

0 ₹ CRN 76-05-1 CMF C2' H F3 02

F- C- CO2H

220046-88-8 CAPLUS
1-Epperazinebutanamide, 4-(1,3-benzodioxol-5-ylmethyl)-β(cyclopentylmethyl)-N-hydroxy-α-[[(4-methoxyphenyl)sulfonyl]amino]γ-οχο-, (αS,βR)-, mono(trifluoroacetate) (salt) (9Cl) Z 3

Erich Leeser

50613257

(CA INDEX NAME)

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CRN 220046-57-1 CMF C29 H38 N4 O8 S

Absolute stereochemistry.

3

CRN 76-05-1 CMF C2 H F3 O2

F-C-C02H

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT REFERENCE COUNT:

14 CAPLUS COPYRIGHT 2007 ACS on STN
1979:604719 CAPLUS
91:204719
Pharmaceutical compositions containing piperazinyl
acylhydroxamic acid derivatives to treat inflammation
or anaphylactic allergy conditions
Coutts, Ronald T.; Biggs, David F.; Wandelmaier, Frank
W.; Semaka, Frank D.
Capadian Patents and Development Ltd., Can.
U.S., 5 pp.
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ΘY ихсоинон Ph (CH2) nN AB Seven piperazinylacylhydroxamic acids I [X = straight or branched C1-3 alkylene, m = 0, 1, or 2, Y = a salt forming acid (when present)] derivs. were prepared by aminoesterification of the corresponding 1-monosubstituted piperazines and then converted to the HC1 salts. The compds. showed antinfiramatory, antianaphylactic, and antidepressant activities. Thus, 2-methyl-3-[1-(4-phenyl)piperazinyl)propionohydroxamic acid-HC1 [71861-77-3] inhibted carrageand-induced edema volume by 23.5% I hafter s.c. administration to rats, decreased egg albumin-induced anaphylaxis by 72% when given i.v. to rats (50 mg/kg), and protected 92% of reserpinized rats given 32 mg of the compound/kg, i.p.
IN 1861-78-44 P 71861-81-9P
RL: SPN (Synthetic preparation)
(preparation and antinflammatory and antianaphylactic activity of)
(preparation and antinflammatory and antianaphylactic activity of)
monohydrochloride (9CI) (CA INDEX NAME) 8

II

S S

CH2-CH-C-NH-OH

● HC1

CH2-CH2-Ph

Z 3

71861-81-9 CAPLUS 1-Piperazinebutanamide, N-hydroxy-4-(2-phenylethyl)-, dihydrochloride (9CI) (CA INDEX NAME)

50613257

(CH2) 3-C-NH-OH CH2-CH2-Ph

●2 HCl

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Erich Leeser